us-10-698-160-1.rge

```
X81785 Epstein-Bar
X81782 Bpstein-Bar
X81786 Epstein-Bar
X81786 Epstein-Bar
BS153 Epstein-Bar
S50728 BNLF! [Epst
U08278 Epstein-Bar
X7442286 Human her
I66494 Seguence 14
BX296528 Home sapi
AC087226 Mus muscu
AC087226 Mus muscu
AC087226 Mus muscu
AC138096 Home sapi
AC079433 Mus muscu
AC079433 Mus muscu
AC079433 Mus muscu
AC07943 Mus muscu
AC07943 Human her
AC18199 Human her
AC18199 Human her
AC180939 Home sapi
AC073554 Home sapi
AC073554 Home sapi
AC073554 Home sapi
AC13809 Macaca mu
AC14308 Macaca mu
AC13808 Macaca mu
AC14308 Macaca mu
                                                                                                                                                                                                                                                                                                                                                             circular VRL 31-JAN-2006
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Farrell, P.J., Bankier, A., Seguin, C., Deininger, P. and Barrell, B.G. Latent and lytic cycle promoters of Epstein-Barr virus EMBO J. 2 (8), 1331-1338 (1983)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Deininger, P.L., Bankier, A., Farrell, P., Baer, R. and Barrell, B. Sequence analysis and in vitro transcription of portions of the Epstein-Barr virus genome
J. Cell. Biochem. 19 (3), 267-274 (1982)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Farrell,P.J., Deininger,P.L., Bankier,A. and Barrell,B.
Homologous upstream sequences near Epstein-Barr virus promoters
Proc. Natl. Acad. Sci. U.S.A. 80 (6), 1565-1569 (1983)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Arrand, J.R., Rymo, L., Walsh, J.E., Bjorck, E., Lindahl, T. and Griffin, B.E.
Molecular cloning of the complete Epstein-Barr virus genome set of overlapping restriction endonuclease fragments
Nucleic Acids Res. 9 (13), 2999-3014 (1981)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Possible role of flanking nucleotides in recognition initiator codon by eukaryotic ribosomes Nucleic Acids Res. 9 (20), 5233-5252 (1981) 7301588
                                                                                                                                                                                                                                                                                                                                                                                                                          Human herpesvirus 4
Viruses; dsDNA viruses, no RNA stage; Herpesviridae;
                                                                                                                                                                                                                                                                                                                                                           HHV507799 171823 bp DNA cir
Human herpesvirus 4 complete wild type genome.
AJ507799
                                                                                                                                                                                                                                                                                                                                                                                                              Human herpesvirus 4 (Epstein-Barr virus)
                                                                                                                                                                                                                                                                                                                                                                                                                                              Gammaherpesvirinae; Lymphocryptovirus
                                                                                                                                                                                                                                                                                                          ALIGNMENTS
                     EBVLMP227
EBVLMP230
                                        HS4LMPA
S50728
EBU08278
HEEBVMP
                                                                                                                 AC087227
AC087226
AC138096
                                                                                                                                                       AC084799
AC079433
AC021347
                                                                                                                                                                                                                      AY519199
AC143808
AC018939
AC073554
                                                                                                                                                                                                  AC094274
AC079425
                                                                                                        BX296528
                                                                                                                                                                                         AC079424
                                                                                              I66494
                                                                                                                                                                                                                                                                                                                                                                                          GI:86261677
  204
204
204
1397
4997
2400
1422
7218
                                                                                                                205691
265985
56970
                                                                                                                                                        303091
252689
                                                                                                                                                                                                                                           134940
185822
253711
                                                                                                                                                                             80133
209887
                                                                                                                                                                                                  235310
222707
                                                                                                                                                                                                                      2431
122272
                                                                                                                                                 497
                                                                                                                                                                                                                                                                                                                                                                                                     complete genome.
                                                                                                                                                                                                                                                                                                                                                                                          AJ507799.2
 Kozak, M.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              6296170
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       6300857
                   199.2
199.2
185.4
                                                                                                     81.6
81.2
81
                                                                                                                                    800.8
800.2
777.4
75.6
75.6
74.8
74.8
                                                                                                                                                                                                                                                                                                                                                           LOCUS
DEFINITION
ACCESSION
VERSION
KEYWORDS
SOURCE
ORGANISM
                                                                                                                                                                                                                                                                                                                                       RESULT 1
HHV507799
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   PUBMED
REFERENCE
AUTHORS
TITLE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         REFERENCE
AUTHORS
TITLE
JOURNAL
PUBMED
REFERENCE
AUTHORS
TITLE
JOURNAL
PUBMED
REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     PUBMED
REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AUTHORS
TITLE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AUTHORS
                                                                                                                                                                                                                                                                                                                                                                                                                                                        REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        JOURNAL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PUBMED
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          JOURNAL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   JOURNAL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      TITLE
  0000000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         V01555 Epstein-Bar
M2066 Epstein-Bar
M2066 Epstein-Bar
X58140 Human herpe
K02165 Epstein-Bar
W66104 Human herpe
K6861 Epstein-Bar
M81720 Epstein-Bar
AJZ78794 Human her
AJZ78794 Human her
AJZ78795 Human her
AJZ78796 Human her
AJZ78796 Human her
AJZ18796 Human her
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AJ507799 Human her
                                                                    Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                    1 tcagggcagtgtgtcaggag.......gcgaaaagggtgtgggctgt 575
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Description
                                                                                                                                                                                                             12732272
         GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd
                                                                                                                                                                                      6366136 segs, 31973710525 residues
                                                                                                                                                                                                       Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               SUMMARIES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    HS4B958RAJ
HS4LMP
AY961628
HBNLF1
HS4LMP1
HS4MP
HHU86104
EBVLMP1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                HHV507799
EBV
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      HS4LMPB
HHE278797
HHE278798
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        HHE278796
EBAJ1676
EBVLMP228
                                                 nucleic search, using sw model
                                                                                                                                                                                                                                                                                    summaries
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      HHE278794
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 HHE278795
                                                                                                                                                      IDENTITY NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                             Minimum Match 0%
Maximum Match 100%
Listing first 45 su
                                                                                                                                                                                                                                length: 0
length: 2000000000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   US-10-698-160-1
575
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Query
Match Length DB
                                                                                                                                                                                                                                                                                                                9b_pat: *
9b_pat: *
9b_pat: *
9b_pr: *
9b_pr: *
9b_er: *
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              100.0 171823
100.0 172281
100.0 172281
90.3 194113
86.6 2038
86.6 2038
84.2 3407
84.2 3407
86.8 2031
74.4 515
74.4 515
74.4 515
70.5 515
70.5 515
70.3 515
64.1 406
                                                                                                                                                                                                                                                                                                        GenEmbl:*
                                                                      June
                                                                                                                                                                                                                                                                                                                                                           Post-processing:
```

Bed 8ed

88

Minimum Maximum

Database

Perfect score:

Sequence:

ı

OM nucleic

Run on:

Scoring table:

Searched:

ø

8

the AUG

ŏ

575 575 575 536.6 504.6 6484.2 464.8 451.2 451.2 428 428 408 405.6 405.6

000000

Score

Result No.

Ab220967 Animal te Aba220967 Animal te Aba622496 Oligonucl Aba652496 Oligonucl Aba652056 Oligonucl Aba652056 Oligonucl Aba652057 Oligonucl Aba652057 Oligonucl Aba682936 Breast ca Aca03973 Human PKD Acu03973 Human Sch Acu03108 Human Sch Acu031281 Cotton an Acu031281 Cotton an Acu031281 Oligonucl Aba035496 Oligonucl Aba03520 Oligonucl Aba036708 Human GPC

ABZ20967
AAA02488
AAA02504
AAA02504
AAA02504
AAA02504
AAA030228
AAA030228
AAA03101
AAT94108
AAT94108
AAT94108
AAT1851
AAT94108
AAT1851
AAT1851
AAT1851
AAT1851
AAT194108
AAT19506
AAC1023807
ABG032495
ABG032495
ABG032495
ABG032495
AAC10130
AAC40130

53522 53522 53522 53526 53526 53577 71131 840 1130 1030 922 1030

500 1593 32195 518 518 1327 100976

```
0000000
                                                                                                                                                                000
                                                           6, 2006, 17:26:38 ; Search time 433.304 Seconds (without alignments) 9252.273 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                         Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                    575
                                                                                                US-10-698-160-1
575
1 tcagggcagtgtgtcaggag......gcgaaaaagggtgtgggctgt
                                                                                                                                                                                 10489840
         GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
                                                                                                                                                                               Total number of hits satisfying chosen parameters:
                                                                                                                                                               5244920 segs, 3486124231 residues
                                                                                                                                                                                                                             Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                           - nucleic search, using sw model
                                                                                                                                     IDENTITY NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                  geneseqn2003bs: *
geneseqn2003cs: *
geneseqn2003ds: *
geneseqn2004as: *
                                                                                                                                                                                                                                                                          geneseqn1990s:*
geneseqn2000s:*
geneseqn2001as:*
geneseqn2001bs:*
geneseqn2001bs:*
geneseqn2001bs:*
geneseqn2002bs:*
                                                                                                                                                                                                                                                                                                                                                                                                geneseqn2005s:*
                                                                                                                                                                                                                                                                                                                                                                                                         geneseqn2006s:*
                                                                                                                                                                                                   Minimum DB seq length: 0
Maximum DB seq length: 200000000
                                                                                                                                                                                                                                                                   N Genesed 8:*
                                                             June
                                                                                                                                                                                                                                                                                                               66...
99...
110...
113...
154...
                                                                                                                                                                                                                               Post-processing:
                                                                                                 Title:
Perfect score:
                                                                                                                                     Scoring table:
                                           OM nucleic
                                                                                                                    Sequence:
                                                                                                                                                                Searched:
                                                                                                                                                                                                                                                                   Database
                                                             Run on:
```

ALIGNMENTS

Score Match Length DB ID Description

575 100.0 1177 12 ADO43140 Ado43140 Epstein-B
575 100.0 172281 12 AD043140 Ad042158 Epstein-B
69.2 12.0 1371 12 AD043140 Ad012158 Epstein-B
69.2 12.0 1371 12 AD012158 Ad012158 Epstein-B
69.2 11.0 1373 10 ADC86736 Adc86736 Human GPC
67.4 11.7 3133 10 ADC86736 Adc86738 Human GPC
66.8 11.6 629 13 ACM54594 Ad086731 Vector pB
66.8 11.6 12733 9 ACD13843 Ad048592 Vector pB
66.8 11.6 12739 9 ACD13843 ACM54596 Ad06868 Human GPC
66.8 11.1 117 10 ADC86688 Add37384 Human GPC
66.8 11.1 2013 9 ACD13843 Ad066688 Human GPC
66.8 11.6 12739 6 ABK98592 Ad046868 Human GPC
66.8 11.1 117 10 ADC86688 Ad06688 Human GPC
67.6 10.9 1065 6 ABT09682 Ad037384 Human CD1
67.6 10.7 28198 10 ADC87789 Ad0377080 Mouse pla
67.6 10.7 28198 10 ADC87789 Ad06778 Human PAL
67.6 10.7 28198 10 ADC87789 Ad06778 Human PAL
67.6 10.7 28198 10 ADC87789 ADC9678 Human PAL

υυ

RESULT 1	1.04.0
	ADO43140 standard; DNA; 1177 BP.
	ADO43140;
17 21 24 24 25 25 25 25 25 25 25 25 25 25 25 25 25	12-AUG-2004 (first entry)
	Epstein-Barr virus LMP1 gene upstream region.
ઇસ લ્લાહ્	<pre>Bpstein-Barr virus; LMPl; promoter; adenovirus; vector; gene therapy; cytostatic; ds:</pre>
	Human herpesvirus 4.
FH	Kev Location/Oualifiers
	tein_bind
PT	/*tag= a /bound_moiety= "STAT"
	•
FT FT	protein_bind 642650 /*tag= b
FT	
	`
	GC_signal 657673 /*tag= c
×	
	WO2004042025-A2.
	21-MAY-2004.
	30-OCT-2003; 2003WO-US034858.
	01-NOV-2002; 2002US-0423203P.
	(CELL-) CELL GENESYS INC.
·	Yu DC, Ramesh N;
DR WP DR GB	WPI; 2004-390609/36. GENRANK; NC 001345.

SUMMARIES

Result No. AG448338 Mus muscu AG134650 Pan trog1 DV792917 Hw\_lotin\_1 DV792917 Hw\_lotin\_1 DV792917 Hw\_lotin\_1 BR733207 FGA507896 B1416480 hasp001xa CR24856 Forward s CR24819 CH261-172 DV7781216 Hw\_Fat\_30 DV7781216 Hw\_Fat\_30 DV7781216 Hw\_Fat\_30 DV775320 Hw\_Fat\_36 CR905417 tric074xf CF805417 tric074xf CF805417 tric074xf CF805417 tric074xf CF805417 tric076xi DV775155 Hw\_Fat\_36 DV775155 Hw\_Fat\_36 DV776929 Hw\_Fat\_36

```
1421 bp mRNA linear EST 28-WAR-2005
CEC06-C08.x1d-t SHGC-CEC Gasterosteus aculeatus cDNA clone
CEC06-C08 3', mENA sequence.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Gasterosteus aculeatus (three spined stickleback)
Gasterosteus aculeatus
Bukaryota, Metazoa, Chordata, Craniata, Vertebrata, Euteleostomi,
Actinopterygii, Metazoa, Theoreti, Euteleostei, Neoteleostei,
Actinopterygii, Percomorpha, Gasterostei, Rectleostei, Neoteleostei,
Acanthomorpha, Acanthopterygii, Percomorpha, Gasterosteiformes,
Gasterosteidae, Gasterosteus.
1 (bases I to 1421)
1 (bases I to 1421)
Schmutz, D. M., Peichel, C., Balabahdra, S., Grimwood, J., Dickson, M.,
Schmutz, J. and Myers, R.M.
Expressed sequence tags from Gasterosteus aculeatus
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Unpublished (2003)
Contact: Grimwood, Jane
Stanford Human Genome Center
Stanford University School of Medicine
975 California Ave, Palo Alto, CA 94304, USA
Tal: 650 320 5901
Fax: 650 320 5801
Email: jane@ehgc.stanford.edu
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 /organism="Gasterosteus aculeatus"
/mol_type="mRNA"
/strain="Conner Creek sticklebacks,
/db xref="taxon:69293"
/clone="CECO6-CO8"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        sex="mixed male and female"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            ALIGNMENTS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 High quality sequence start: 4
High quality sequence stop: 347
Location/Qualifiers
                                                                DR733207
BH416480
CB905439
CF876953
CR234856
CC281819
DV775320
DV775320
DV775320
CC8905417
CF876937
CF876937
CF876937
CF876937
CF876937
CF876937
CF876937
CF876937
DV794756
DV794756
DV794756
                                                                                                                                                                                                                                                                                                                                                                                  AG448781
CL078355
CG754612
DN710897
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        tissue_type="skin"
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                DN653570.1 GI:61958819
                                                                                                                                                                                                                                                                                           01111111
                                                                                                                                       422244
                                                                                                                                                                                                                                           1013
1087
1087
1487
666
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  1. .1421
                                                                                                   1142
1142
680
1040
1449
1525
805
                                                                                                                                                                                                                                                                                                                                                1320
1335
11578
11598
11793
                                                   133.9
133.9
133.9
133.9
133.9
133.9
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
133.7
BST
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              VERSION
KEYWORDS
SOURCE
ORGANISM
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    source
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            DEFINITION
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       REFERENCE
AUTHORS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     TITLE
JOURNAL
COMMENT
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              ACCESSION
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            RESULT 1
DN653570
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  FEATURES
O
                                                                                                                                                                                                           O
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        DV779627 Hw Pat 35
CL045106 CH216-62E
AG36566 Mus muscu
CL081488 CH216-162
CC255263 CH264-42A
BQ441712 AGENCOURT
CW948789 TCB33.1 H
BE040676 OF04404 Ō
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 CL077232 CH216-143
CG750956 P045-2-H0
CW951178 TCB36.3 D
BM556092 AGENCOURT
DV781271 Hw Fat 35
CL509275 SAIL 811
CL082769 CH216-171
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           N653570 CEC06-C08
                                                                                                              6, 2006, 17:52:13; Search time 3888.44 Seconds (without alignments) 8269.032 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           N808203 74134031
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                                                                  US-10-698-160-1
575
1 tcagggcagtgtgtcaggag......gcgaaaagggtgtgggctgt 575
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Description
                                                                                                                                                                                                                                                                                                                                                96473596
              GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
                                                                                                                                                                                                                                                                                                           48236798 segs, 27959665780 residues
                                                                                                                                                                                                                                                                                                                                           Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         SUMMARIES
                                                                                                                                                                                                                                                                                                                                                                                                                                 Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                              OM nucleic - nucleic search, using sw model
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DN808203
DV779627
CL045106
AG363665
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               CL081488
CC255263
BQ441712
CW948789
BC40676
CB905469
CF876976
CL077232
CC0750956
CW551178
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           DN653570
                                                                                                                                                                                                                                                       IDENTITY NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                                               seq length: 0
seq length: 200000000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             14
12
13
13
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Length DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     9b_est1:*
9b_est3:*
9b_est5:*
9b_est6:*
9b_est7:*
9b_est9:*
9b_est9:*
9b_gss1:*
9b_gss1:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         1607
1129
1301
1560
1593
1021
1287
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 849
1591
1738
956
1362
1471
1099
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        EST: *
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query
Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           44444444
                                                                                                                   June
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              10:
                                                                                                                                                                                                                                                                                                                                                                                                                                  Post-processing:
                                                                                                                                                                                                    Perfect score:
                                                                                                                                                                                                                                                       Scoring table:
```

0B 0B

Minimum

Database

Searched:

Sequence:

Run on:

/dev stage="adult" /lab\_host="DH10B (Tl phage resistant)" /clone\_lib="SHGC-CEC" /note="Vector: Express 1; Total and poly A+ RNA was

DV781271 CL509275 CL082769

υv

8.11.8 8.11.6

טט

υυ

Score 82

Result No.

WA USA"

```
Sequence 14, Appl
Sequence 4735, Ap
Sequence 2, Appli
Sequence 1, Appli
Sequence 16801, A
Sequence 12147, A
Sequence 17361, A
Sequence 13443, A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              11, Appl
4508, Ap
14, Appl
558, App
                                                                                                                   June 6, 2006, 18:05:44; Search time 153.504 Seconds (without alignments) 7008.836 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   1880ed_Patents_NA:*
1: /EMC_Celerra_SIDS3/ptodata/2/ina/5_COMB.seq:*
2: /EMC_Celerra_SIDS3/ptodata/2/ina/5_COMB.seq:*
3: /EMC_Celerra_SIDS3/ptodata/2/ina/6A_COMB.seq:*
4: /EMC_Celerra_SIDS3/ptodata/2/ina/6B_COMB.seq:*
5: /EMC_Celerra_SIDS3/ptodata/2/ina/H_COMB.seq:*
6: /EMC_Celerra_SIDS3/ptodata/2/ina/H_COMB.seq:*
7: /EMC_Celerra_SIDS3/ptodata/2/ina/PCTUS_COMB.seq:*
8: /EMC_Celerra_SIDS3/ptodata/2/ina/PCTUS_COMB.seq:*
9: /EMC_Celerra_SIDS3/ptodata/2/ina/PCTUS_COMB.seq:*
10: /EMC_Celerra_SIDS3/ptodata/2/ina/PCTUS_COMB.seq:*
                                                                                                                                                                                                                                           1 tcagggcagtgtgtcaggag......gcgaaaagggtgtgggctgt 575
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Description
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Sequence Seq
GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
                                                                                                                                                                                                                                                                                                                                                                                       2807332
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     US-08-232-463-14
US-08-628-136-2
US-08-658-136-2
US-08-658-136-1
US-09-949-016-12147
US-09-949-016-12147
US-09-949-016-12147
US-09-949-016-1343
US-09-949-016-1343
US-09-949-016-1343
US-09-949-016-1318
US-09-949-016-1318
US-09-949-016-1318
US-09-165-264-14
US-09-165-264-14
US-08-123-98-558
US-08-123-98-558
US-08-123-98-558
US-08-123-98-558
US-08-99-34-958
US-09-99-34-958
US-09-99-34-058
US-09-99-34-058
US-09-007-005-17
US-09-244-796-17
                                                                                                                                                                                                                                                                                                                                                                               Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                             1403666 segs, 935554401 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    SUMMARIES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                           OM nucleic - nucleic search, using sw model
                                                                                                                                                                                                                                                                                IDENTITY NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                                                                                           seq length: 0
seq length: 200000000
                                                                                                                                                                                                 US-10-698-160-1
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Length DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Query
                                                                                                                                                                                                                      Perfect score:
                                                                                                                                                                                                                                                                                  Scoring table:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Minimum DB
Maximum DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         Database :
                                                                                                                                                                                                                                             Sequence:
                                                                                                                       Run on:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Result
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Š.
```

Sequence 16945, A Sequence 12, Appli Sequence 12, Appli Sequence 3, Appli Sequence 11042, A Sequence 11042, A Sequence 12403, A Sequence 12160, A Sequence 12160, A Sequence 1726, A Sequence 317, Appli Sequence 317, Appli Sequence 317, Appli Sequence 3175, Appli Sequence 321, Appli Sequence 323, Appli Sequence 328, Appli Sequence 328, Appli Sequence 4415, Appli Sequence 4415, Appli Sequence 4415, Appli Sequence 4724,		
24 49.8 8.7 37155 3 US-09-949-016-16945 25 49.6 8.6 319 3 US-09-165-264-8 26 48.6 8.5 320 3 US-09-165-264-12 27 48.6 8.5 4897 10 5196516-7 29 48 8.3 10968 3 US-09-165-264-7 29 47 8.2 12907 3 US-09-249-016-16505 23 47 8.2 12907 3 US-09-949-016-16505 23 47 8.2 119032 3 US-09-949-016-12403 24 47 8.2 119032 3 US-09-949-016-12160 25 4 47 8.2 15331 3 US-09-949-016-1260 25 4 47 8.2 155331 3 US-09-949-016-1260 26 35 46.2 8.0 1235 4 US-09-949-016-1260 27 4 45.8 8.0 1235 4 US-09-949-016-1260 28 45.8 8.0 1235 4 US-09-949-016-1260 29 45.8 10 1017 4 US-09-297-648-4525 20 45 45.2 7.9 1166 3 US-09-037-297-297-328 20 44 44.8 7.8 1411 4 US-09-297-648-415	RESULT 1  US-08-212-463-14/C  Sequence 14, Application US/08232463  PRETAL NO. 5670367  GENERAL INFORMATION: PRACTORY: APPLICANT: SCHEIFLINGER, F. APPLICANT: SCHEIFLINGER, F. APPLICANT: SCHEIFLINGER, F. APPLICANT: PALKNER, F. G. TITLE OF INVENTION: RECOMBINANT FOWLPOX VIRUS NUMBER OF SEQUENCES: 52 CORRESPONDENCE ADDRESS: STATE: 1800 Diagonal Road, Suite 500 CITY: Alexandria STREFT: 1800 Diagonal Road, Suite 500 CITY: Alexandria STREET: 1800 Diagonal Road, Suite 500 CITY: Alexandria STREET: 1800 Diagonal Road, Suite 500 CITY: Alexandria STREET: 1800 Diagonal Road, Suite 500 COMPUTER: USA COUNTER: USA COUNTER: USA COMPUTER: IBM PC Compatible OPERATING SYSTEM: PC-DOS/MS-DOS SOFTWARE: Patentin Release #1.0, Version #1.25 COMPUTER: 1800 PC COMPATION MEDIUCATION NUMBER: US/08/232,463 FILING DATE: CLASSIFICATION NUMBER: US/07/935,313 FILING DATE: APPLICATION NUMBER: EP 91 114 300.6 FILING DATE: APPLICATION NUMBER: BP 91 114 300.6 FILING DATE: APPLICATION NUMBER: BP 91 114 300.6 FILING DATE: APPLICATION NUMBER: 30472/114 IMMU TELERRICE CHARACTERISTICS: APPLICATION NUMBER: 30472/114 IMMU TELERRICE CHARACTERISTICS: LENGTH: 7218 Dass Pairs INFORMATION FOR SEQ ID NO: 14: SECONETIC CHARACTERISTICS: LENGTH: 7218 Dass Pairs TTELENGTH: 7218 Dass Pairs TTELENG	; TOPOLOGY: linear ; IMMEDIATE SOURCE:

```
Published Applications NA Main:*

1: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US07_PUBCOMB.seq:*

2: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US08_PUBCOMB.seq:*

3: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US09_PUBCOMB.seq:*

4: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US09_PUBCOMB.seq:*

5: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*

7: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*

8: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*

9: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*

10: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*

11: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*

12: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*

13: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10G_PUBCOMB.seq:*

14: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*

15: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*

16: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*

16: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*

16: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*
                                                                                                                                                                                                       6, 2006, 18:14:24 ; Search time 1119.71 Seconds (without alignments) 6310.019 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                     1 tcagggcagtgtgtcaggag......gcgaaaagggtgtggggctgt 575
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         37784340
GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            18892170 seqs, 6143817638 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                                                                                     OM nucleic - nucleic search, using sw model
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 IDENTITY_NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Minimum DB seq length: 0
Maximum DB seq length: 200000000
                                                                                                                                                                                                                                                                                                                                                                   US-10-698-160-1
                                                                                                                                                                                                                      June
                                                                                                                                                                                                                                                                                                                                                                                                      Perfect score:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Scoring table:
                                                                                                                                                                                                                                                                                                                                                                                                                                            Sequence:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Searched:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Database
                                                                                                                                                                                                                  Run on:
```

SUMMARIES	Description	JS-10-698-160-1 Sequence 1, Appli	Seguence		Seguence			US-10-779-543-5579 Sequence 5579, Ap	Ø			Sequence	Sequence	-	Sequence	Segmence	Sequence	
	рв ір	. 6	0 6	8	8	8	9	10	7	7	9	7	7	7	7	9	9	
	Query Match Length	575	891	925	778	815	985	1337	5452	5452	166	3163	3163	3133	3133	629	12733	
dp	Query Match	100.0	12.6	12.6	12.3	12.1	12.1	12.0	11.9	11.9	11.8	11.8	11.8	11.7	11.7	11.6	11.6	
	Score	575	72.4	72.2	70.6	9.69	69.4	69.2	68.2	68.2	67.8	67.6	67.6	67.4	67.4	66.8	66.8	
	Result No.		7 0	ი ი	4	C 52	9	0	80	6	c 10	11	12	13	14	15	16	

B US-10-437-963-28390 Sequence B US-10-425-115-14077 Sequence B US-10-021-323-3377 Sequence C US-10-021-323-3377 Sequence C US-10-022-798-1141 Sequence C US-0-292-798-1141 Sequence D US-09-804-682-33 Sequence D US-10-425-115-11145 Sequence D US-10-425-115-11404 Sequence D US-10-425-115-26028 Sequence D US-10-425-13908 Sequence D US-10-425-3908 Sequence D US-10-425-3908 Sequence D US-10-363-4834-3908 Sequence D US-10-363-4834-3908 Sequence D US-10-425-135-146990 Sequence D US-10-425-15-146990 Sequence D US-10-425-115-146990 Sequence D US-10-425-115-14690	υ	18	66.4	11.5	822	σ	US-10-425-115-102047	Sequence 102047,
11.2   700 9 US-10-425-115-14077   Sequence   11.1   588 8 US-10-021-33-9377   Sequence   11.1   1117 7 US-10-021-33-9377   Sequence   11.1   1117 7 US-10-021-33-9377   Sequence   10.9   1065 3 US-09-804-682-33   Sequence   10.9   1065 3 US-09-804-682-33   Sequence   10.7   771 8 US-10-425-115-9386   Sequence   10.7   771 8 US-10-425-115-9386   Sequence   10.6   1064 3 US-10-425-115-9386   Sequence   10.6   804 8 US-10-425-115-9386   Sequence   10.5   692 9 US-10-425-115-26028   Sequence   10.6   600 9 US-10-363-345A-39087   Sequence   10.3   600 9 US-10-363-3483A-39087   Sequence   10.3   600 9 US-10-363-3483A-39089   Sequence   10.3   600 10 US-10-363-3483A-39089   Sequence   10.1   959 8 US-10-425-995-90899   Sequence   10.0   674 8 US-10-437-963-64501   Sequence   10.0   087 00 00 00 00 00 00 00 00 00 00 00 00 00	υ	13	65.2	11.3	862	ω	839	
11.1   588	O	20	64.2		700	σ		14077
11.1   1117 7 US-10-1017-161-1403   Sequence   10.9   10.55 3 US-09-084-682-3   Sequence   10.9   10.55 3 US-09-084-682-3   Sequence   10.7   771   8 US-10-424-599-107958   Sequence   10.7   771   8 US-10-425-115-93385   Sequence   10.6   673   9 US-10-425-115-93385   Sequence   10.6   673   9 US-10-425-115-93385   Sequence   10.6   673   9 US-10-425-115-93385   Sequence   10.5   604   3 US-10-425-115-1145   Sequence   10.5   600   8 US-10-425-115-10535   Sequence   10.5   600   8 US-10-425-115-6258   Sequence   10.5   600   8 US-10-424-599-100535   Sequence   10.5   600   9 US-10-424-599-81404   Sequence   10.3   600   9 US-10-424-599-81404   Sequence   10.3   600   0 US-10-363-445-3908   Sequence   10.3   600   0 US-10-363-445-3908   Sequence   10.3   600   0 US-10-363-445-3908   Sequence   10.1   2432   8 US-10-437-963-64501   Sequence   10.0   677   9 US-10-437-963-64501   Sequence   10.0   677   9 US-10-425-115-146990   Sequence   10.0   0 US-0000171159A1		21	63.6		588	89	US-10-021-323-9377	
11.1   1117 7 US-10-292-799-1141   Sequence   10.8   10.65 3 US-0-804-682-33   Sequence   10.8   10.65 3 US-0-804-682-33   Sequence   10.7   1211 9 US-10-425-115-93385   Sequence   10.7   1211 9 US-10-425-115-13385   Sequence   10.6   10.64 3 US-10-425-115-13385   Sequence   10.6   10.64 3 US-10-425-115-1145   Sequence   10.6   10.64 3 US-10-425-115-26028   Sequence   10.5   692 9 US-10-425-115-26028   Sequence   10.5   692 9 US-10-425-115-26028   Sequence   10.5   692 9 US-10-425-115-26028   Sequence   10.3   600 9 US-10-425-13-8575   Sequence   10.3   600 9 US-10-425-13-9087   Sequence   10.3   600 9 US-10-363-483A-3908   Sequence   10.3   600 10 US-10-363-483A-3908   Sequence   10.3   600 10 US-10-363-483A-3908   Sequence   10.1   959 8 US-10-437-963-64501   Sequence   10.1   959 8 US-10-437-963-64501   Sequence   10.0   674 8 US-10-425-115-146990   Sequence   10.0   0.0		22	63.6		1117	7	US-10-017-161-1403	1403,
10.9 1065 3 US-09-804-682-33 Sequence 10.7 771 8 US-10-779-533-5571 Sequence 10.7 771 8 US-10-425-115-93365 Sequence 10.6 673 US-10-425-115-11145 Sequence 10.6 673 US-09-804-682-29 Sequence 10.6 804 8 US-10-424-59-100535 Sequence 10.5 500 8 US-10-68-7434-2 Sequence 10.5 500 8 US-10-68-7434-2 Sequence 10.5 1087 8 US-10-425-115-26028 Sequence 10.3 600 9 US-10-425-135-26028 Sequence 10.3 600 9 US-10-437-3963-56258 Sequence 10.3 600 9 US-10-437-3963-56258 Sequence 10.3 600 9 US-10-363-34334-39089 Sequence 10.3 600 10 US-10-363-4834-39089 Sequence 10.1 959 8 US-10-779-543-8591 Sequence 10.1 959 8 US-10-779-543-8591 Sequence 10.1 959 8 US-10-779-63-64501 Sequence 10.0 674 8 US-10-779-63-64501 Sequence 10.0 674 8 US-10-779-63-64501 Sequence 10.0 974 8 US-10-779-63-64501 Sequence 10.0 974 8 US-10-779-63-6868 Sequence 10.0 074 8 US-10-779-63-6869 Sequence 10.0 074 8 US-10-779-63-6869 Sequence 10.0 074 8 US-10-75-115-16990 Sequence 10.0 08-10-115-115-115-115-115-115-115-115-115-		23	63.6	11.1	1117	7	US-10-292-798-1141	1141,
10.8 1000 10 US-10-779-543-8571 Sequence 10.7 771 8 US-10-424-599-107958 Sequence 10.7 771 8 US-10-425-115-93385 Sequence 10.6 673 9 US-10-425-115-11145 Sequence 10.6 673 9 US-10-425-115-11145 Sequence 10.5 500 8 US-10-424-599-100535 Sequence 10.5 500 8 US-10-424-599-100535 Sequence 10.5 500 8 US-10-424-599-100535 Sequence 10.5 682 9 US-10-425-115-2628 Sequence 10.5 1087 US-10-424-599-81404 Sequence 10.3 600 9 US-10-424-599-81404 Sequence 10.3 600 9 US-10-424-599-81404 Sequence 10.3 600 9 US-10-343-45A-39088 Sequence 10.3 600 10 US-10-363-345A-39089 Sequence 10.3 600 10 US-10-363-345A-39089 Sequence 10.1 2432 8 US-10-437-963-64501 Sequence 10.0 847 9 US-10-437-963-64501 Sequence 10.0 847 9 US-10-424-599-69089 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 10.0 WS-10-424-599-69089 Sequence 10.0 WS-10-425-115-146990 Sequence 10.0 WS-10-424-599-69089 Sequence 10.0 WS-10-425-115-146990 Sequence 10.0 WS-10-424-599-69089 Se	υ	24	62.6		1065	m	US-09-804-682-33	
10.7 771 8 US-10-424-599-107958 Sequence 10.6 73 9 US-10-425-115-93385 Sequence 10.6 73 9 US-10-425-115-93385 Sequence 10.6 1064 3 US-10-425-115-1145 Sequence 10.6 1064 3 US-10-425-125-1055 Sequence 10.5 692 9 US-10-425-15-26028 Sequence 10.5 692 9 US-10-425-15-26028 Sequence 10.5 1087 8 US-10-425-15-26028 Sequence 10.3 1218 10 US-10-425-135-399-81404 Sequence 10.3 600 9 US-10-363-3457-39087 Sequence 10.3 600 9 US-10-363-3457-39087 Sequence 10.3 600 10 US-10-363-3483A-39087 Sequence 10.3 600 10 US-10-363-3483A-39087 Sequence 10.2 1593 10 US-10-363-4834-3908 Sequence 10.1 959 8 US-10-437-963-64501 Sequence 10.0 674 8 US-10-437-963-64501 Sequence 10.0 674 9 US-10-437-963-64501 Sequence 10.0 877 9 US-10-437-963-64501 Sequence 10.0 877 9 US-10-437-963-64501 Sequence 10.0 877 9 US-10-425-115-146990 Sequence 10.0 877 9 US-10-427-963-64501 Sequence 10.0 WS-10-437-963-64501 Seq		25	62		1000	10	US-10-779-543-8571	
10.7 1211 9 US-10-425-115-93385 Sequence 10.6 673 US-09-804-682-29 Sequence 10.6 804 8 US-09-804-682-29 Sequence 10.5 804 8 US-09-804-682-29 Sequence 10.5 800 8 US-10-424-59-10035 Sequence 10.5 500 8 US-10-425-115-26028 Sequence 10.5 1087 8 US-10-424-59-81404 Sequence 10.3 600 9 US-10-437-963-56258 Sequence 10.3 600 9 US-10-363-345A-39089 Sequence 10.3 600 9 US-10-363-345A-39089 Sequence 10.3 600 10 US-10-363-48591 Sequence 10.1 559 8 US-10-739-543-8591 Sequence 10.1 559 8 US-10-739-543-8694 Sequence 10.1 559 8 US-10-739-543-8694 Sequence 10.1 559 8 US-10-437-963-64501 Sequence 10.0 674 8 US-10-437-963-64501 Sequence 10.0 974 BUS-10-425-115-146990 Sequence 10.0 974 BUS-10-425-115-146990 Sequence 10.0 0 US-10-425-115-146990 Sequence 10.0 0 US-10-425-115-14690 Sequence 10.0 0 US-10-4		56	61.8		771	8	US-10-424-599-107958	Sequence 107958,
10.6 673 9 US-10-425-115-11145 Sequence 10.6 1064 3 US-09-804-682-29 10.6 1064 3 US-09-804-682-29 10.5 500 8 US-10-424-599-100535 Sequence 10.5 692 9 US-10-425-115-26028 Sequence 10.5 692 9 US-10-425-115-26028 Sequence 10.4 545 8 US-10-437-963-56258 Sequence 10.3 600 9 US-10-73-943-3908 Sequence 10.3 600 9 US-10-73-345A-3908 Sequence 10.3 600 10 US-10-363-345A-3908 Sequence 10.3 600 10 US-10-363-345A-3908 Sequence 10.1 5432 8 US-10-437-963-6650 10.1 2432 8 US-10-437-963-64501 Sequence 10.0 847 9 US-10-437-963-64501 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 10.0 847 9 US-10-425-115-16908 Sequence 10.0 WS-10-425-115-16688 Sequence 10.0 WS-10-425-115-16908 Sequence		27	61.4		1211	6	US-10-425-115-93385	Sequence 93385, A
10.6 1064 3 US-09-804-689-29 10.6 804 8 US-10-424-599-100535 Sequence 10.5 500 8 US-10-666-749A-0 10.5 692 9 US-10-666-749A-0 10.4 545 8 US-10-425-115-26028 10.3 600 9 US-10-779-543-8575 10.3 600 9 US-10-779-543-8575 10.3 600 9 US-10-779-543-8575 10.3 600 10 US-10-363-483A-39087 Sequence 10.4 550 10 US-10-363-483A-39087 Sequence 10.5 1081 8 US-10-363-483A-39087 Sequence 10.6 10 US-10-363-483A-39089 Sequence 10.1 2432 8 US-10-363-483A-39089 Sequence 10.1 2432 8 US-10-437-963-64501 Sequence 10.0 674 8 US-10-437-963-64501 Sequence 10.0 674 9 US-10-437-963-68694 Sequence 10.0 877 9 US-10-437-963-68696 Sequence 10.0 878 US-10-437-963-68696 Sequence	υ	28	61		673	σ	US-10-425-115-11145	Seguence 11145, A
10.6 804 8 US-10-424-599-100535 Sequence 10.5 500 8 US-10-668-7949.2 10.5 500 9 US-10-668-799-100535 10.5 1087 8 US-10-425-115-26028 10.4 545 8 US-10-437-595-81404 10.3 600 9 US-10-363-3457-39089 10.3 600 9 US-10-363-3457-39089 10.3 600 10 US-10-363-3457-39089 10.1 529 8 US-10-363-3483-39089 10.1 0.2 1593 10 US-10-363-48391 10.1 0.2 1593 10 US-10-799-543-8591 10.1 0.2 1593 10 US-10-799-6989 10.0 674 8 US-10-437-966-64501 10.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0 0.0	υ	53	61	10.6	1064	٣	US-09-804-682-29	
10.5 500 8 US-10-668-749A-2 10.5 692 9 US-10-425-115-26028 10.6 10.7 10.8 9 US-10-425-115-26028 10.3 1218 10 US-10-425-19404 10.3 1218 10 US-10-779-543-8575 10.3 600 9 US-10-363-345A-39087 10.3 600 10 US-10-363-345A-39088 10.2 1593 10 US-10-363-345A-39088 10.1 2432 8 US-10-363-483A-39088 10.1 2432 8 US-10-363-483A-39088 10.1 2432 8 US-10-437-963-64501 10.1 847 9 US-10-437-963-64501 10.0 847 9 US-10-425-115-146990 10.0 847 9 US-10-425-115-146990 10.0 847 9 US-10-425-115-146990 10.0 US-10-425-115-15-146990		30	ö	10.6	804	80	US-10-424-599-100535	
10.5 692 9 US-10-425-115-26028 Sequence 10.5 1087 8 US-10-424-29-81404 Sequence 10.4 545 8 US-10-427-963-56258 Sequence 10.3 1218 10 US-10-779-543-8575 Sequence 10.3 600 9 US-10-363-345A-39087 Sequence 10.3 600 10 US-10-363-345A-39088 Sequence 10.3 600 10 US-10-363-345A-39088 Sequence 10.1 2432 8 US-10-363-483A-39089 Sequence 10.1 2432 8 US-10-437-963-64501 Sequence 10.0 674 8 US-10-424-599-69089 Sequence 10.0 674 8 US-10-424-599-69089 Sequence 10.0 874 9 US-10-425-115-146990 Sequence 9.9 32195 3 US-10-425-115-146990 Sequence 9.9 32195 3 US-10-425-115-146990 Sequence 9.0 US20040171159A1	U	31			200	8	US-10-668-749A-2	
10.5 1087 8 US-10-424-599-81404 Sequence 10.3 1218 10 US-10-437-959-81409 10.3 600 9 US-10-437-959-81408 Sequence 10.3 600 9 US-10-363-345A-39088 Sequence 10.3 600 10 US-10-363-483A-39088 Sequence 10.3 600 10 US-10-363-483A-39088 Sequence 10.1 10.2 1593 10 US-10-363-4839 Sequence 10.1 2432 8 US-10-363-3899 Sequence 10.1 959 8 US-10-437-963-64501 Sequence 10.0 674 8 US-10-437-963-6868 Sequence 10.0 877 8 US-10-437-963-6868 Sequence 10.0 877 8 US-10-437-963-6868 Sequence 10.0 9.9 32195 3 US-09-764-891-6868 Sequence 9.9 32195 3 US-09-764-891-6868 Sequence 9.0 US20040171159A1	O	32			692	0	US-10-425-115-26028	26028,
10.4 545 8 US-10-437-963-55258 Sequence 5 10.3 1218 10 US-10-779-543-8575 Sequence 5 10.3 600 9 US-10-7363-345A-39087 Sequence 3 10.3 600 9 US-10-363-345A-39088 Sequence 10.3 600 10 US-10-363-483A-39088 Sequence 10.3 600 10 US-10-363-483A-39088 Sequence 10.2 1593 10 US-10-779-543-8591 Sequence 10.1 2432 8 US-10-779-543-8591 Sequence 6 10.0 674 8 US-10-779-543-8591 Sequence 6 10.0 674 8 US-10-424-599-6089 Sequence 10.0 674 8 US-10-425-115-146990 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 6 9.9 32195 3 US-09-764-891-6868 Sequence 6 0.0 US20040171159A1	υ	33	60.2		1087	8	US-10-424-599-81404	
10.3 1218 10 US-10-779-543-8575 Sequence 10.3 600 9 US-10-363-345A-39087 Sequence 10.3 600 9 US-10-363-345A-39089 Sequence 10.3 600 10 US-10-363-483A-39088 Sequence 10.1 591 0 US-10-363-483A-39088 Sequence 10.1 12432 8 US-10-775-543-891 Sequence 10.1 2432 8 US-10-437-963-64501 Sequence 10.0 674 8 US-10-437-963-64699 Sequence 10.0 674 8 US-10-424-599-69089 Sequence 6 9.9 32195 3 US-09-764-891-6868 Sequence 6 9.9 32195 3 US-09-764-891-6868 Sequence 6 0.0 US20040171159A1	Ü	34	59.6	10.4	545	œ	US-10-437-963-56258	56258,
10.3 600 9 US-10-363-345A-19087 Sequence 3 10.3 600 10 US-10-363-485A-39088 Sequence 1 10.3 600 10 US-10-363-483A-39089 Sequence 1 10.3 600 10 US-10-363-483A-39089 Sequence 1 10.1 2432 8 US-10-779-543-6591 Sequence 1 10.1 259 8 US-10-437-963-64501 Sequence 6 10.0 674 8 US-10-437-963-69089 Sequence 6 10.0 874 9 US-10-424-5999 Sequence 6 10.0 874 9 US-10-424-5990 Sequence 6 10.0 874 9 US-10-424-5990 Sequence 6 10.0 875 10-10-16688 Sequence 6 10		35	59.4	10.3	1218	10	US-10-779-543-8575	e 8575,
10.3 600 9 US-10-363-345A-1908B Sequence 3 600 10 US-10-363-485A-19087 Sequence 10.3 600 10 US-10-363-485A-19087 Sequence 10.2 1593 10 US-10-363-48591 Sequence 10.1 2432 8 US-10-779-543-8591 Sequence 10.1 2432 8 US-10-437-963-64501 Sequence 6 10.0 674 8 US-10-424-599-69089 Sequence 10.0 847 9 US-10-424-599-69089 Sequence 10.0 847 9 US-10-424-115-146990 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 10.0 B47 9 US-10-424-115-146990 Sequence 10.0 B47 9 US-10-424-115-146990 Sequence 10.0 US-10-424-115-14690 Sequence 10.0 US-10-424-115-115-14690 Sequence 10.0 US-10-424-115-115-14690 Sequence 10.0 US-10-424-115-115-115-115-115-115-115-115-115-11		36	59.2	10.3	009	σ	US-10-363-345A-39087	39087,
10.3 600 10 US-10-363-483A-39087 Sequence 10.1 593 10 US-10-363-483A-39088 Sequence 10.2 1593 10 US-10-779-543-8591 Sequence 10.1 2432 8 US-10-779-543-8659 Sequence 10.1 959 8 US-10-437-963-64501 Sequence 10.0 674 8 US-10-425-99-6089 Sequence 10.0 847 9 US-10-424-599-6089 Sequence 10.0 847 9 US-10-424-115-146990 Sequence 10.0 847 9 US-10-425-115-1668 Sequence 10.0 B47 9 US-10-6458 Sequence 10.0 B47 9 US-10-6458 Sequence 10.0 US-10-6458 Sequence 10.0 US-10-6458 Sequence 10.0 US-10-6458 Sequence 10.0 US-10-698160 Sequence	U	37	59.2	10.3	600	0	US-10-363-345A-39088	39088,
10.3 600 10 US-10-36-483A-3908B Sequence 10.1 1593 10 US-10-779-543-8591 Sequence 10.1 2432 8 US-10-779-543-8591 Sequence 10.1 959 8 US-10-437-963-64694 Sequence 10.0 674 8 US-10-424-599-69089 Sequence 10.0 847 9 US-10-424-5990 Sequence 9.9 32195 3 US-09-764-891-6868 Sequence 6 Dilication US/10698160 0. US20040171159A1		38	59.2	10.3	900	10	US-10-363-483A-39087	
10.2 1593 10 US-10-779-543-8591 Sequence 10.1 2432 8 US-10-437-963-64501 Sequence 10.0 674 8 US-10-437-963-64594 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 9.9 32195 3 US-09-764-891-6868 Sequence Philoation US/10698160 o. US20040171159A1	υ	33	59.2	10.3	009	10	US-10-363-483A-39088	
10.1 2432 8 US-10-437-963-64501 Sequence 6 10.1 959 8 US-10-437-963-48694 Sequence 4 10.0 674 8 US-10-424-599-65089 Sequence 1 10.0 847 9 US-10-425-115-146990 Sequence 1 9.9 32195 3 US-09-764-891-6868 Sequence 6 9.9 32195 3 US-09-764-891-6868 Sequence 6 Pplication US/10698160 O. US20040171159A1		40	58.4	10.2	1593	10	US-10-779-543-8591	
10.1 959 8 US-10-437-963-48694 Sequence 4 10.0 674 8 US-10-424-599-69089 Sequence 6 10.0 847 9 US-10-424-599 Sequence 6 9.9 32195 3 US-09-764-891-6868 Sequence 6 ALIGNMENTS  PDlication US/10698160 O. US20040171159A1	O	41	58	10.1	2432	æ	US-10-437-963-64501	
10.0 674 8 US-10-424-599-69089 Sequence 10.0 847 9 US-10-425-115-146990 Sequence 9.9 32195 3 US-09-764-891-6868 Sequence ALIGNMENTS  PPlication US/10698160 0. US20040171159A1 MATTON:	O	42	۲.	10.1	959	œ	-437-963-4869	4
10.0 847 9 US-10-425-115-146990 Sequence 9.9 32195 3 US-09-764-891-6868 Sequence ALIGNMENTS  PDlication US/10698160 o. US20040171159A1 MATION:		43	۲.	10.0	674	œ	-424-599-6908	69069
9.9 32195 3 US-09-764-891-6868 Sequence ALIGNMENTS  PD11cation US/10698160 o. US20040171159A1 MATION:	υ	44	۲.	10.0	847	σ	-425-115-14699	146990
pplication US/1069816 o. US20040171159A1 MATION:	U	45	ė.	6.6	0	9	-09-764-891-686	
RESULT 1 US-10-698-160-1 ; Sequence 1, Application US/10698160 ; Publication No. US20040171159A1 ; GENERAL INFORMATION:							ALIGNMENTS	
US-10-698-160-1 ; Sequence 1, Application US/10698160 ; Publication No. US20040171159A1 ; GENERAL INFORMATION:	RES	SULT 1	_					
; Sequence 1, Application US/10698160 ; Publication No. US20040171159A1 ; GENERAL INFORMATION:	as	-10-69	1-091-8					
GENERAL INFORMATION:		Seguen	ation N	pplicat	ion US/	1065	18160	
A DISTRIBUTION OF US		FINERA	I INFOR	MATTON	4 4 4 4 5 4 5 4 5 6 6 6 6 6 6 6 6 6 6 6	2		
		A DDT.T	CANT	Ş				

120 180 240 61 GGCTTATGTAGGGCGGCTACGTCAGAGTAACGCGTGTTTCTTGGGATGTAGGCCCGGGGG 120 121 GAITTGCGGGGTCTGCCGGAGGCAGTACGGGTACAGATTTCCCGAAAGCGGCGGTGTGTG 180 9 9 1 TCAGGGCAGTGTGAGGAGCAAGCAGTTGAGGAAAGAAGGGGGCCAGAGCAGTGTGAGA TCAGGGCAGTGTGTCAGGAGCAAGTTGAGGAAAGAAGAAGGGGCCAGAGCAGTGTGAGA 61 GCCTTATGTAGGCGCCGCTACGTCAGAGTAACGCGTGTTTCTTGGGATGTAGGCCCGGGGG TGTGCATGTAAGCGTAGAAAGGGGAAGTAGAAAGCGTGTGTTTGTGTTAGAAAAGCGGGT 121 GATTTGCGGGGTCTGCCGGAGGCAGTACGGGTACAGATTTCCCGAAAGCGGCGGGGTGTGTG Gaps ô Length 575; 100.0%; Score 575; DB 8; Length 57 100.0%; Pred. No. 5.1e-163; tive 0; Mismatches 0; Indels APPLICANT: U. 14
APPLICANT: U. 14
APPLICANT: U. 14
TITLE OF INVENTION: CELL-SPECIFIC ADENOVIRUS VECTOR
TITLE OF INVENTION: COMPRISING EBV-SPECIFIC PROMOTER
FILE REFERENCE: CELL-0.3
CURRENT APPLICATION NUMBER: US/10/698,160
CURRENT FILING DATE: 2003-10-30
PRIOR PILING DATE: 2002-11-01
NUMBER OF SEQ ID NOS: 4
SOFTWARE: PastSEQ for Windows Version 4.0
SEQ ID NO 1
LENGTH 575 ; TYPE: DNA; ORGANISM: Epstein Barr virus US-10-698-160-1 Query Match Best Local Similarity 100.0 Matches 575; Conservative н 181 ð 윱 ò g ò 요 8

ö

```
June 6, 2006, 18:40:59; Search time 37:9911 Seconds (without alignments) 1782.525 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Published Applications NA New.*

1. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US09_NEW_PUB.seq:*

2. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US06_NEW_PUB.seq:*

3. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US07_NEW_PUB.seq:*

4. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US08_NEW_PUB.seq:*

5. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US08_NEW_PUB.seq:*

7. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10_NEW_PUB.seq:*

7. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10_NEW_PUB.seq:*

7. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10_NEW_PUB.seq:*

7. /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10_NEW_PUB.seq:*
                                                                                                                                                                                                                                                                                                                                                                 1 tcagggcagtgtgtcaggag.......gcgaaaagggtgtgggctgt 575
GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               tal number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       246837 segs, 58886990 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Dst-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                                                                          - nucleic search, using sw model
                                                                                                                                                                                                                                                                                                                                                                                                                            IDENTITY_NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              inimum DB seq length: 0
                                                                                                                                                                                                                                                                                                      US-10-698-160-1
                                                                                                                                                                                                                                                                                                      Title:
Perfect score:
                                                                                                                                                                                                                                                                                                                                                                                                                         Scoring table:
                                                                                                                          OM nucleic
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             atabase :
                                                                                                                                                                                                                                                                                                                                                                    Sequence:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          arched:
                                                                                                                                                                                  Run on:
```

#### 23294, A 38422, A 1885, Ap 1885, Ap 16, Appl 32064, A 1481, Ap 1481, Ap 37022, A 32708, A 1477, Ap 29733, A 63, Appl 28, Appl Sequence 31197, A Description Sequence US-10-519-335-37 US-10-953-349-35485 US-11-293-697-14477 US-11-293-697-14477 US-11-293-697-1476 US-11-297-134-697 US-11-293-697-1476 US-11-293-697-1496 US-11-293-697-1885 US-10-953-349-25482 US-11-284-877-16 US-11-284-877-16 US-11-284-877-16 US-11-293-697-1881 US-11-293-697-1881 US-11-293-697-1881 US-11-294-877-16 US-11-294-877-16 US-11-294-877-16 US-11-294-877-16 US-10-953-349-31614 SUMMARIES 8 Query Match Length 1116 1683 1799 464 22118 1773 38187 128361 151830 435 1214 2244 1951 11978 37476 4004 3029 Score 33.8 Result

Sequence 31008, A Sequence 38692, A Sequence 3879, A Sequence 3579, A Sequence 3572, A Sequence 31282, A Sequence 390, App Sequence 1893, App GENERAL INFORMATI Sequence 318, A Sequence 23388, A Sequence 1797, App Sequence 21797, App Sequence 21797, App Sequence 21797, App Sequence 2165, App Sequence 516, App Sequence 516, App Sequence 2165, App Sequence	ULT 1   10-953-349-31197/c   2010953349   2010953349   2010953349   2010953349   2010953349   2010953349   2010953349   20109503451   201095030050107345A1   201095030050107345A1   201095030050107345A1   201095030050107345A1   201095030050107345A1   201095030050107345A1   201095030050107345A1   2010950300501073745A1   201095030050107349   201	AGACGCGGA 522            TGACGTGGAA 262 GALIFOTNIA Stic Kidney Disease in the Cat
US-10-953-349-31008 US-10-953-349-38692 US-10-953-349-35793 US-10-953-349-35793 US-10-953-349-35793 US-11-293-697-390 US-11-293-697-390 US-11-293-697-390 US-10-953-349-3581 US-10-953-349-3338 US-10-953-349-3338 US-10-953-349-3338 US-11-293-697-611 US-11-293-697-611 US-11-293-349-21053 US-10-953-349-21053 US-10-953-349-21053 US-10-953-349-21053 US-10-953-349-21053	US/10953349 5A1 colai et al. E-DETERMINED DNA FRAGME ED THERBY US/10/953,349 09-30 09-30 3.3 mm	AGAGGGCTAAGGGTCGGCGGCAGCAGCAGCAGCCGGAA GGCGGGCGTGTTGTACTCGGTGTTGACGAAGCTGCTGGAA  Application US/10857260 No. US20060110742A1 RWATION: Lyons, Leslie A. Grahn, Robert Erdman, Robert Erdman, Carolyn The Regents of the University of California VEMTION: Carrier Tests for Polycystic Kidney NCE: 021070-146800US LICATION NUMBER: US/10/857,260
32 2 5 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6	ULT 1  10-953-349-31197/c  equence 31197, Application US/10  UBLICALION NO. US20060107345A1  ENERAL INFORMATION:  APPLICANT: ALEXANDROV, NICKOLAI  TITLE OF INVENTION: ENCOUNED DITTLE OF INVENTION: ENCOUNED THATLE OF INVENTION SEQ ID NOS: 40252  SOFTWARE: PARCHING DATE: 2004-09-30  NUMBER OF SEQ ID NOS: 40252  SOFTWARE: PARCHINI VERSION 3.3  LENGTH: 1682  TYPE: DNA  10-953-349-31197  10-953-349-31197  10-953-349-31197  424 AGAGGGGGGGGCAACC  360 AGAGGGGGGGGCCAACC  424 AGAGCGCGGGGATGTCCTGCC  424 AGAGCGCAGGGATGTCTGCCAACC  10 MILL OF MACHINIAN STANCE OF ACCORDANCE OF ACCORDANC	480 AGAGGGTAAGGGTCGGCGGGTGA
	RESULT 1 US-10-953-349-31197/c ; Sequence 31197, App; ; Publication No. US2; ; GENERAL INFORMATION ; TITLE OF INVERTION ; TITLE OF INVERTION ; TITLE OF INVERTION ; CURRENT FILIGO DATE ; SOFTWARE: PAPELICATION ; SOFTWARE: PAPELICATION ; SOFTWARE: PAPENILD ; TYPE: DNA ; TYPE: DNA ; ORGANISM: TITLICU US-10-953-349-31197 OUETY MATCH BEST LOCAL SIMILIARI MATCHES 87; CONS  QY 360 AGAGGGC  QY 424 AGAGGGC  QY 420 GGGGGGGT  DD 364 TGGTGGAG	Oy 480 AGAGG Db 304 GGCGG RESULT 2 US-10-857-260-31/c ; Sequence 31, Appl ; Publication No. U ; GENERAL INFORMATI ; APPLICANT: LYONE ; APPLICANT: Grah ; APPLICANT: Grah ; APPLICANT: The ; TITLE OF INVENTI ; FILE REFERENCE: ; CURRENT APPLICANT: ; FULE REFERENCE: ; CURRENT APPLICANT: ; FULE REFERENCE: ; CURRENT APPLICANT: ; FULE REFERENCE:

Medical T Sequence

score:

Perfect

Sequence:

ŧ

OM nucleic

Run on:

Scoring table:

Searched:

Database

```
AX743545 Sequence
AX766903 Sequence
AX766903 Sequence
AX766903 Sequence
AX766903 Sequence
AX766903 Sequence
AX766903 Sequence
AX766909 Sequence
166494 Sequence
AC077494 Homo sapi
AC077494 Homo sapi
AC0719146 Homo sapi
AC019146 Homo sapi
AC019146 Reattus no
AC118496 Reattus no
AC119677 Peppio anu
AC093831 Homo sapi
AC093831 Homo sapi
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       circular VRL 31-JAN-2006
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          Farrell, P.J., Bankier, A., Seguin, C., Deininger, P. and Barrell, B.G.
Latent and lytic cycle promoters of Epstein-Barr virus
EMBO J. 2 (8), 1331-1338 (1983)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Deininger, P.L., Bankier, A., Farrell, P., Baer, R. and Barrell, B. Sequence analysis and in vitro transcription of portions of the Epstein-Barr virus genome
J. Cell. Biochem. 19 (3), 267-274 (1982)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Parrell, P.J., Deininger, P.L., Bankier, A. and Barrell, B.
Homologous upstream sequences near Epstein-Barr virus promoters
Proc. Natl. Acad. Sci. U.S.A. 80 (6), 1565-1569 (1983)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Griffin, B.E. Molecular cloning of the complete Epstein-Barr virus genome set of overlapping restriction endonuclease fragments Nucleic Acids Res. 9 (13), 2999-3014 (1981)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    Possible role of flanking nucleotides in recognition of initiator codon by eukaryotic ribosomes Nucleic Acids Res. 9 (20), 5233-5252 (1981)
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Arrand, J.R., Rymo, L., Walsh, J.E., Bjorck, E., Lindahl, T.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Human herpesvirus 4
Viruses; dsDNA viruses, no RNA stage; Herpesviridae;
Gammaherpesvirinae; Lymphocryptovirus.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          инуэи7799 171823 bp DNA сіл
Human herpesvirus 4 complete wild type genome.
AJ507799
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 complete genome.
Human herpesvirus 4 (Epstein-Barr virus)
                                                                                                                                                                                                                                                                                                                                                      LMFL5213T
CT005267_08
AC119057
                                                                                                                                                                                                                                                                                                                                                                                                                                                                         ALIGNMENTS
                                                                                                                                                                                                                                                                                           CT005271_13
CP000096_09
AC118496
                                                                                                                                                                                                           AC079429
AC027494
AC091905
                                                                                                                                                                                                                                                             AC019146
AK026479
                               AX743545
AX766903
AX774657
DD211187
AX743546
AX766958
AX7655393
AX655393
AX655393
AX6007950
AC0023591
                                                                                                                                                                                                                                                                                                                                                                                                                        AC120554
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    AJ507799.2 GI:86261677
                                                                                                                                                                                                                                                                                                                                                         112
                                                                                                                                                                                                                                            2 12 2
                                                                                                                                                                                                                                                                                         122
50
50
50
50
50
1250
1250
170979
                                                                                                                                                                                            171456
258174
                                                                                                                                                                                                                                                                                                                                                                                       190427
182116
165083
                                                                                                                                                                                                                                           158741
189592
                                                                                                                                                                                                                                                                                                                         241138
                                                                                                                                                                                                                               156987
                                                                                                                                                                                                                                                                                                          110000
 00.0000000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        0872327
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Cozak, M.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             6296170
37.8
37.6
37.6
37.6
37.4
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 DEFINITION
ACCESSION
VERSION
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                KEYWORDS
SOURCE
ORGANISM
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      RESULT 1
HHV507799
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              REFERENCE
AUTHORS
TITLE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            PUBMED
AUTHORS
TITLE
JOURNAL
PUBMED
AUTHORS
AUTHORS
AUTHORS
AUTHORS
AUTHORS
TITLE
JOURNAL
PUBMED
PUB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              PUBMED
REFERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              AUTHORS
TITLE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                REPERENCE
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  AUTHORS
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 PUBMED
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                JOURNAL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             JOURNAL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             JOURNAL
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  TITLE
 LOCUS
                                                                                 0000
                                                                                                                                                           0 0000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         AJ507799 Human her V0155 Epstein-Bar AY961628 Human her AJ414532 Human her AJ414534 Human her AJ414534 Human her AJ414535 Human her AJ414535 Human her AJ414535 Human her AJ414535 Homo sapi AK27652 Homo sapi AK27652 Bpstein-Bar AX930675 Sequence CQ77455 Sequence CQ77455 Sequence CQ881165 Sequence CQ881165 Sequence CQ881165 Sequence CQ881165 Sequence
                                                                                                       6, 2006, 17:51:08 ; Search time 3285.57 Seconds (without alignments) 10607.392 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             results predicted by chance to have a l to the score of the result being printed, of the total score distribution.
                                                                                                                                                                                                        egeaccaactgeggeaaatg.........cattteaggeeggttttgea 545
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Description
                                                                                                                                                                                                                                                                                                                        12732272
             GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
                                                                                                                                                                                                                                                                                      6366136 segs, 31973710525 residues
                                                                                                                                                                                                                                                                                                                 Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              SUMMARIES
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AY961628
HHE414532
HHE414534
HHE414534
HHE414535
CQ850504
AX03785
AX03785
AX127652
AX127652
AX127652
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          HS4B958RAJ
                                                                                                                                                                                                                                                                                                                                                                                                   Minimum Match O%
Maximum Match 100%
Listing first 45 summaries
                                                                           nucleic search, using sw model
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           HHV507799
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        AX930675
CQ774555
CQ794832
CQ881165
CQ891130
                                                                                                                                                                                                                                       IDENTITY_NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             Pred. No. is the number of res
score greater than or equal to
and is derived by analysis of
                                                                                                                                                                                                                                                                                                                                                    Minimum DB seq length: 0
Maximum DB seq length: 200000000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              US-10-698-160-2
545
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                              Length DB
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 9b_env:*
9b_pat:*
9b_pat:*
9b_pi:*
9b_ro:*
9b_ro:*
9b_ov:*
9b_nv:*
9b_nv:*
9b_nv:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       100.0 171823
100.0 172281
100.0 1842113
99.7 171657
75.4 426
772.9 425
72.9 17096
13.8 2035
13.8 2035
13.8 2035
13.8 2035
8.3 50
8.3 50
8.3 50
                                                                                                                                                                                                                                                                                                                                                                                                                                                                  GenEmbl:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            Query
Match
                                                                                                          June
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                66:
77:
77:
110:
113:
114:
                                                                                                                                                                                                                                                                                                                                                                                                    Post-processing:
```

ø

ав

the AUG

Score

Š Result

545 545 545

543.4 410.8 4410.8 397.2 397.2 110.6 75

111111111 116543216987654321

50.4

```
6, 2006, 17:26:38 ; Search time 410.696 Seconds (without alignments) 9252.273 Million cell updates/sec
                                                                                                                                                                                                                             1 cgcaccaactgcggcaaatg......satttcaggccggttttgca 545
                                                                                                                                                                                                                                                                                                                                                              10489840
GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
                                                                                                                                                                                                                                                                                                                                                          Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Ą.
                                                                                                                                                                                                                                                                                                                         5244920 segs, 3486124231 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                        Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                            - nucleic search, using sw model
                                                                                                                                                                                                                                                                IDENTITY NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               geneseqn2003cs:*geneseqn2003ds:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                    geneseqn2004as:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      geneseqn2004bs:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     geneseqn2002as:*geneseqn2002bs:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        geneseqn2003as:*
geneseqn2003bs:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         geneseqn2001as:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 geneseqn2001bs:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        geneseqn2005s:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         geneseqn2006s:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   geneseqn1980s:*
geneseqn1990s:*
                                                                                                                                                                                                                                                                                                                                                                                                    Minimum DB seq length: 0
Maximum DB seq length: 2000000000
                                                                                                                                                                                        US-10-698-160-2
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     N Geneseq 8:*
                                                                                                                 June
                                                                                                                                                                                                         Perfect score:
                                                                                                                                                                                                                                                                  Scoring table:
                                                                            OM nucleic
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Database :
                                                                                                                                                                                                                               Sequence:
                                                                                                                                                                                                                                                                                                                         Searched:
                                                                                                                 Run on:
```

SUMMARIES

	Description	Ado43141 Epstein-B	Adn12161 Epstein-B	Adt05874 Cercopith	Adr07467 Full leng	Abz77478 Oligonucl	Acc83339 TP1 promo	Abz81622 TP1 promo	Acc84643 TP1 promo	Acc78948 TP1 promo	Acf58279 Oligo use	Adh39824 TPl promo	Adj45801 TP1 promo	Adl73003 Notch Big	Adg14415 TP1 promo	Ad852104 Notch sig	Acc83340 TP1 promo	Acf58280 Oligo use	Ad173004 Notch sig
	A .	.2 ADO43141	2 ADN12161	.3 ADT05874	3 ADR07467	1 ABZ77478	1 ACC83339	ABZ81622	ACC84643	ACC78948	.0 ACF58279	.0 ADH39824	.2 ADJ45801	.2 ADL/3003	.2 ADQ14415	.3 ADS52104	3 ACC83340	.0 ACF58280	.2 ADL73004
	Match Length DB ID	545 1	172281 1	171096 1	2035 1	S0 8	50 B	50 8	20	20	50	50	50 1	50	50	50	20	50	50
Query	Match	1.00.0	100.0	20.3	13.8	8.3	8.3	8.3	8.3	8.3	8.3	8.3	8.3	8.3	8.3	8.3	8.1	8.1	8.1
	Score	545	545	110.6	75	45.4	45.4	45.4	45.4	45.4	45.4	45.4	45.4	45.4	45.4	45.4	44	44	44
Result	No.	1	~	m	4	'n	9	7	60	o	10	11	12	13	14	15	c 16	c 17	c 18

Ada71938 Rice gene Aas85600 DNA encod	Adz64267 Human can Adz64254 Human can		Marker		AeeU5181 Cancer-as		σ	Abl29468 Drosophil		Abk62959 Rat seque	Aah18665 Human cDN	Aak84206 Human imm	Aas32548 Human gen	Aas32546 Human gen	Adb58672 Toxicity-	Adc26226 Human NOV	Aas41696 Genomic s	Aba20646 Human ner	Adq59368 Human can		Add71350 Glutamine	Human	Acl26294 Rice abio	
ADA71938 AAS85600	ADZ64267 ADZ64254	AAL53530	ADJ75856	ADZ60295	AEE05181	AEB32388	ADS36459	ABL29468	ABL77620	ABK62959	AAH18665	AAK84206	AAS32548	AAS32546	ADB58672	ADC26226	AAS41696	ABA20646	ADQ59368	ADL13497	ADD71350	ADQ21602	ACL26294	
യഹ	14	9	77	7	4.	14	13	4	9	9	4	4	4	4	10	20	4	Ŋ	12	ព	10	77	7	
2000	5791 15363	3635	3635	3635	156296	160300	165156	4786	351	502	5008	7177	7471	8319	14746	849	9481	9481	100944	91823	93390	131673	1965	
7.6	6.8 6.8	9.9	9.9			9.9	6.5	6.5	6.5	6.4	6.4	6.4	6.4	6.4	6.4	6.2	6.2	6.2	6.2	•	6.2	•	6.1	
41.4	37	36.2	36.2	36.2	9 0		35.6	35.4	35.2	34.8	34.8	34.8	34.8	34.8	34.8	34	33.8	33.8	33.8	33.6	33.6	33.6	33.4	
c 20	21 22	c 23	C 24		50	28	29	30	31	c 32	c 33	m	35	36	37	c 38	39	40	41	c 42	43	44	45	

## ALIGNMENTS

RESULT 1 AD043141

Epstein-Barr virus; LMP2A; promoter; adenovirus; vector; gene therapy; Epstein-Barr virus LMP2A promoter. ADO43141 standard; DNA; 545 BP. (first entry) Human herpesvirus 4. WO2004042025-A2. cytostatic; ds. 12-AUG-2004 AD043141;

30-OCT-2003; 2003WO-US034858. 01-NOV-2002; 2002US-0423203P. 21-MAY-2004. 

(CELL-) CELL GENESYS INC.

WPI; 2004-390609/36. Ramesh N; Yu DC,

New replication-competent adenovirus vector comprising an adenovirus gene essential for replication, useful as a vehicle for introducing new genetic capability. Claim 7; SEQ ID NO 2; 36pp; English.

The present sequence is the Epstein-Barr virus (EBV) LMP2A gene promoter region. The LMP2 gene encodes 2 distinct proteins, LMP2A and LMP2B. LMP2A aggregates in patches within the plasma membrane of latently infected B-lymphocytes. The consistent expression of LMP2A in Hodgkin's disease and nasopharyngeal carcinoma suggests an important function for this protein in oncogenes. The invention provides replication-competent adenovirus

```
June 6, 2006, 17:52:13; Search time 3685.56 Seconds (without alignments) 8269.032 Million cell updates/sec
                                                                                                                                                       US-10-698-160-2
545
1 cgcaccaactgcggcaaatg......catttcaggccggttttgca 545
                                                                                                                                                                                                                                                                                                       96473596
GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
                                                                                                                                                                                                                                                                   48236798 seqs, 27959665780 residues
                                                                                                                                                                                                                                                                                                  Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                                                               4
                                                                                                                                                                                                                                                                                                                                                                               Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                           OM nucleic - nucleic search, using sw model
                                                                                                                                                                                                                       IDENTITY NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                Minimum DB seq length: 0
Maximum DB seq length: 200000000
                                                                                                                                                                                                                                                                                                                                                                                                                                                         9b est1::
9b est2::
9b est5::
9b est6::
9b est6::
9b est7::
9b est7::
9b est7::
9b est8::
9b est8::
9b est8::
9b est8::
                                                                                                                                                                                                                                                                                                                                                                                                                                               EST: *
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            10:
                                                                                                                                                       Title:
Perfect score:
                                                                                                                                                                                                                       Scoring table:
                                                                                                                                                                                                                                                                                                                                                                                                                                               Database :
                                                                                                                                                                                      Sequence:
                                                                                                                                                                                                                                                                     Searched:
                                                                                           Run on:
```

	Description	DA645119 DA645119	DA644493 DA644493	CA472883 AGENCOURT	AL059400 Drosophil	AG106932 Pan trodl	DT890143 1471125 M	_	CE082035 tigr-gss-	AQ531217 RPCI-11-3	CB852782 UI-CF-FN0	AG536872 Mus muscu	DR112573 RTS1 29 D	AL050923 Drogophil	BJ306689 BJ306689	AL107369 Drosophil	AG898391 Oryza sat	DT891511 1472661 M	AQ389958 RPCI11-15	AL106358 Drosophil
SUMMARIES	ID	DA645119	DA644493	CA472883	CNSOOCNG	AG106932	DT890143	CC268659	CE082035	AQ531217	CB852782	AG536872	DR112573	CNS007WH	BJ306689	CNS016YN	AG898391	DT891511	AQ389958	CNS0166K
	B	6	0	4	14	14	20	12	12	11	4	14	σ	14	~	14	14	10	11	14
	Watch Length DB	587	524	1359	939	823	780	1079	375	588	294	650	661	692	405	1101	996	784	352	1225
ع	Match	13.8	0.6	8.1	8.0	7.8	7.5	7.4	7.3	7.3	7.2	7.2	7.2	7.2	7.2	7.2	7.1	7.1	7.0	7.0
	Score	75	<b>4</b> ,	44	43.8	42.4	40.8	40.4	39.8	39.6	39.4	39.4	39.4	39.4	39.5	39.5	38.8	38.6	38.4	38
1,000	No.	1	63	e U	Ω	ល	9	0	80	σ υ	10	c 11	12	c 13	c 14	c 15	c 16	c 17	c 18	c 19

/organism="Homo sapiens" /mol\_type="mRNA"

17.8 6.9 401 4 BY284357 17.6 6.9 114 4 12 CC258918 17.4 6.9 114 4 12 CC258918 17.4 6.9 729 5 CD871322 17.2 6.8 779 4 13 CW668045 17.2 6.8 696 10 DV371960 17.2 6.8 698 11 AZ007222 17.2 6.8 950 11 AZ007222 17.2 6.8 950 11 AZ007222 17.2 6.8 950 11 AZ007222 17.3 6.8 1267 12 CC26090 18.6 6.7 728 14 DX304408 18.7 6.6 628 14 GNS0658N 18.2 6.6 628 14 BY28669 18.2 6.6 628 14 BY28669 18.2 6.6 628 14 BY28669 18.2 6.6 628 1 BB847659 18.3 7 8 BY292031 18.4 6.7 728 14 DX304408 18.4 6.7 728 14 DX304408 18.4 6.7 728 14 DX304408 18.4 6.7 728 14 BY28669 18.4 6.7 728 14 BY28669 18.4 6.7 728 14 BY28669 18.4 6.7 728 14 BY28699 18.4 6.7 728 14 BY28999 18.4 14 CNORDER 1 BR1114 EREPTORTER 1 BR11 RETERTOR 0 TECHER 1 BY	BY284357 BY284357 DN370251 LIB3733-0 CC58918 CH361-164	BY424295 BY424295	CW668045 OG_BBa002	CB854072 UI-CF-DUI CF891453 UI-CF-FN0	DV317960 NABOBSITF AZO07202 RDCT-23-3	DB333210 DB333210	ALUSEBSZ DIOBODILL CB591225 AGENCOURT	CC260090 CH261-165	AL065838 Drosophil AL052823 Drosophil	DX304808 OR ABA021	CC184353 CH261-61H	BY293974 BY293974	BY292031 BY292031	BY296096 BY296096	B1288669 B1288669 BB656179 BB666170	BB641431 BB641431 BY728949 BY728949				mRNA linear EST 08-NOV-2005	2,				iata; Vertebrata; Buteleostomi; es; Primates; Catarrhini;		Ota,T., Nishikawa,T.,	sono, Y., Irie, R., Kushida, N.,	Yokoi,T., Kondo,H., Wagatsuma,M., I., Takahashi-Fujii,A	akai, K., Isogai, T. and Sugano, S.	Modulation: Large-scale n of Putative Alternative					Chiba, 292-0818, Japan			and Industrial Technology	; cunA library construction: -end one pass sequencing: HRI,	logy (RAB) and Biotechnology			
ที่ที่ที่ที่ที่ที่ที่ที่ที่ที่ที่ที่ที่ท	BY284357 DN370251	BY424295	CW66804	CF891453	DV317960 1 AZ007202	DB333210	CB591225	2 CC260090	CNS004M4	1 DX304808	2 CC184353	BY293974 BB847650	BY292031	BY296096	B1288669 BB656178	BB641431 BY728949	ALIGNMENTS			587 bp	sapiens cDNA	46			Chordata; Cran Euarchontoglir		I, A., Suzuki, Y.	Saito, K., I	R., Kanda,K., S., Ishibashi,	Kikuchi, H., N	ranscriptional haracterizatio	enes	(9007) 69-66	ਜੂ ੀ	m) tute	i, Kisarazu, Cl		y.com	ect (New Energy	zation, Japan) tute (HRI); 5'	for Biotechnol	יייינית מד ופכוח	alifiers	8m="Homo sapiens"
ที่ที่ที่ที่ที่ที่ที่ที่ที่ที่ที่ที่ที่ท	401 554 1144	411	797	969	968 508	539	904	1267	588 9 80	728	1193	345	387	392	430	667				ou de la constant de	ьтмрви ношс		ens (human)	00	; Metazoa; Eutheria;	; nome: 1 to 587)	, Wakamatsu	Sugiyama, 7	T., Otsuka, K., Ishida,	, Nagai, K.,	cation of T ation and C	of Human	1/11 01 .8	Takao Isoga	ct (AKI Tea earch Insti	ива-Катаtаr	38-52-3975 38-52-3986	j-cdna@nift	n cDNA proj	ncal Organi earch Insti	Association atfonal Inc	encing: RAB	Location/Qu	1587 /organism="
	7.8 6.7.8	7.4	. 4. C	7.2 6.	7.2 6.	80		8.0	6.4.	.4	.4	2.0		. 6	9.0	9 9						DA645119 DA645119.		Homo sapi	Eukaryota Mammalia; Hominidae	1 (bases	Kimura, K.	Ishii, S.,	Yoneyama, Murakawa,	Tanase, T.	Diversiti Identific	Promoters	16344560	Contact:	Helix Res	2-6-7 Kaz	Tel: 81-4	Email: fl	NEDO huma	Helix Res	Research	pass segu		
EDD ESSENS E O E	20	53	52.4			30		33	3 5	36	37	20 G 70 C	6 4	41	1. 4 4. 6.	4.4		RESULT 1	DA645119	LOCUS	DEFINITION	ACCESSION	SOURCE	ORGANISM		REFERENCE	AUTHORS			1	TITLE	TAMBLE	를 급	COMMENT									FEATURES	sonice

Appli 

```
32
24
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                15440, A
14182, A
4514, Ap
144470,
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       15203, A
15204, A
252, App
12787, A
12043, A
15436, A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          17249, A
13317, A
15197, A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Sequence 14, Appl
Sequence 6, Appli
                                                                                                                                                 June 6, 2006, 18:05:44; Search time 145.496 Seconds (without alignments) 7008.836 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      14950, A
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.
                                                                                                                                                                                                                                                                                              545
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Description
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           /EMC_Celerra_SIDS3/ptodata/2/ina/backfiles1.seg:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          BENC_Celerra_SIDS3/ptodata/2/ina/1_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/5_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/6_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/6_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/6_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/H_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/H_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/PCTUS_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/PCTUS_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/PCTUS_COMB.seq:*
| EMC_Celerra_SIDS3/ptodata/2/ina/PCTUS_COMB.seq:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            seduence Sed
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Sequence
Sequence
Sequence
Sequence
Sequence
                                                                                                                                                                                                                                             US-10-698-160-2
545
1 cgcaccaactgcggcaaatg......cattcaggccggtttgca
                       5.1.9
Biocceleration Ltd.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                             US-08-232-463-14

US-09-349-016-12319

US-09-949-016-12319

US-09-949-016-128658

US-09-949-016-11840

US-09-949-016-11840

US-09-949-016-11289

US-09-949-016-11249

US-09-949-016-11217

US-09-949-016-11219

US-09-949-016-118219

US-09-949-016-118219

US-09-949-016-118218

US-09-949-016-118218

US-09-949-016-118218

US-09-949-016-118218

US-09-949-016-118218

US-09-949-016-118218

US-09-949-016-118218

US-09-949-016-118203

US-09-949-016-118203
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  -09-949-016-144470
                                                                                                                                                                                                                                                                                                                                                                                                                                                  Total number of hits satisfying chosen parameters:
                                                                                                                                                                                                                                                                                                                                                                                                          1403666 segs, 935554401 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 SUMMARIES
                       GenCore version (c) 1993 - 2006
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                                                            - nucleic search, using sw model
                                                                                                                                                                                                                                                                                                                                        IDENTITY NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Issued Patents NA:*
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Minimum DB seq length: 0
Maximum DB seq length: 2000000000
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       8
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          24984
63860
99304
422592
1185
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 15004
11366
44377
44378
35337
40548
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  601
11280
11280
43991
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       Length
                                          Copyright
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Ouery
Match
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Title:
Perfect score:
Sequence:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                               Scoring table:
                                                                                                              OM nucleic
                                                                                                                                                                                                                                                                                                                                                                                                               Searched:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Database
                                                                                                                                                        Run on:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                Result
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ģ
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           0 0 0
```

Sequence

44554 44555 199945

Sequence Seq		
C 24 32.2 5.9 421118 3 US-09-949-016-16297 26 32 3.9 123463 3 US-09-949-016-17078 27 31.8 5.8 601 3 US-09-949-016-13840 28 31.8 5.8 601 3 US-09-949-016-54264 29 31.6 5.8 1256 2 US-08-176-4278-9 2 31 31.6 5.8 1256 3 US-08-674-5098-5 2 32 31.6 5.8 1256 3 US-08-674-5098-5 2 35 31.6 5.8 1256 3 US-08-674-5098-5 2 36 31.6 5.8 1256 3 US-08-674-5098-5 2 37 31.6 5.8 1256 3 US-09-639-689-5 2 38 31.6 5.8 1256 3 US-09-639-695-5 2 39 31.6 5.8 1256 3 US-09-448-188-5 2 39 31.6 5.8 1256 3 US-09-448-188-5 2 40 31.6 5.8 1256 3 US-09-448-188-5 2 41 31.6 5.8 1256 3 US-09-448-188-5 2 42 31.6 5.8 1256 3 US-09-418-128-5 2 43 31.6 5.8 1256 3 US-09-418-28-5 2 44 31.6 5.8 1256 3 US-09-536-476-5 2 45 31.6 5.8 1256 3 US-09-136-476-5 2 47 31.6 5.8 1256 3 US-09-131-3-31-3-31-3-31-3-3-31-3-3-3-3-3-3-	RESULT 1 US-08-232-463-14/C 1 Sequence 14, Application US/08232463 2 Sequence 14, Application US/08232463 3 Sequence 14, Application US/08232463 3 Sequence 14, Application US/08232463 5 Sequence 14, Application US/08232463 5 SEQUENCE 1000 6 TITLE OF INVENTION: RECOMBINANT FOWLPOX VIRUS NUMBER OF SEQUENCES: 52 CORRESPONDENCE ADDRESS: 52 CORRESPONDENCE ADDRESS: 52 CORRESPONDENCE ADDRESS: 52 CORRESPONDENCE ADDRESS: 1800 Diagonal Road, Suite 500 CITY: Alexandria STATE: VA COUNTRY: USA ZIP: 22313-029 COMPUTER READABLE FORM: MEDIUM TYPE: IBM PC COMPALING SOFTWARE: Patentin Release #1.0, Version #1.25 CURRENT APPLICATION NUMBER: US/08/232,463 FILING DATE: FILING FI	PRIOR APPLICATION DATA. APPLICATION NUMBER: US/07/935,313 FILING DATE. APPLICATION NUMBER: US/07/935,313 FILING DATE: 26-AUG-1991 ATTORNEY/AGENT INFORMATION: NAME: BENT, Stephen A. REGISTRATION NUMBER: 29,768 REFERENCE/DOCKET NUMBER: 30472/114 IMMU TELECOMMUNICATION INFORMATION: TELECOMMUNICATION INFORMATION: TELEFAX: (703)83-4109

us-10-698-160-2.rnpbm

```
Published Applications NA Main:*

1: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US07_PUBCOMB.seq:*
2: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US08A_PUBCOMB.seq:*
3: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US08A_PUBCOMB.seq:*
4: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US09B_PUBCOMB.seq:*
5: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10A_PUBCOMB.seq:*
6: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
7: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
8: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
9: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
10: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
11: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
12: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
13: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US10B_PUBCOMB.seq:*
14: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*
15: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*
16: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*
16: /EMC_Celerra_SIDS3/ptodata/2/pubpna/US11B_PUBCOMB.seq:*
                                                                                                                                                                                  6, 2006, 18:14:24; Search time 1061.29 Seconds (without alignments) 6310.019 Million cell updates/sec
                                                                                                                                                                                                                                                                                                                                                                                  1 cgcaccaactgcggcaaatg......satttcaggccggttttgca 545
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          37784340
5.1.9
Biocceleration Ltd.
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           18892170 segs, 6143817638 residues
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                   Total number of hits satisfying chosen parameters:
   GenCore version Copyright (c) 1993 - 2006
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Post-processing: Minimum Match 0%
Maximum Match 100%
Listing first 45 summaries
                                                                                                                       - nucleic search, using sw model
                                                                                                                                                                                                                                                                                                                                                                                                                                           IDENTITY NUC
Gapop 10.0 , Gapext 1.0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     seq length: 0
seq length: 200000000
                                                                                                                                                                                                                                                                                                                US-10-698-160-2
545
                                                                                                                                                                                         June
                                                                                                                                                                                                                                                                                                                Title:
Perfect score:
                                                                                                                                                                                                                                                                                                                                                                                                                                              Scoring table:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Minimum DB Maximum DB M
                                                                                                                          OM nucleic
                                                                                                                                                                                                                                                                                                                                                                               Sequence:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           Searched:
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                      Database
                                                                                                                                                                                         Run on:
```

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

Description	Sequence 2, Appli Sequence 180450,	Sequence 180450, Sequence 8, Appli			Sequence 68, Appl Sequence 11, Appl	12,	Sequence 22, Appl Sequence 46, Appl		Sequence 23, Appl	Sequence 6, Appli	Sequence 35, Appl	Semience 12 Anni
SUMMARIES	US-10-698-160-2 US-10-027-632-180450	US-10-027-632-180450 US-10-612-869-8	US-10-763-362-22 US-10-812-144-5	US-10-765-727-34	US-10-846-989-68 US-10-764-415B-11	US-10-845-834A-12	US-10-958-784-22 US-11-078-735-46	US-11-050-346-44	US-10-763-362-23	US-10-812-144-6	US-10-765-727-35	US-10-846-989-12
80	<b>6</b> 40	<b>ر</b> م	ი ი	0	6 2	10	15	16	σ	σ	o	0
* Query Match Length DB	545 612	612 171096	50 00	20	0 0 0 0	20	200	20	20	20	20	20
* Query Match	100.0	86.9 20.3	80 80 61 61	8.3	ໝ ໝ ຕຸຕ	8.9	 	8.3	8.1	8.1	8.1	8.1
Score	545 473.6	473.6	45.4 45.4	45.4	45.4	45.4	45.4	45.4	44	44	44	44
ult No.	- 7	w 4	0 v	7	<b>ω</b> σν	91	11	ជ	7	12	16	17
Result No.	υ	υ							υ	U	Ų	U

ט	18	44	8.1	20	15	US-11-078-735-47	Sequence 47, Appl
υ	13	44	8.1	20	16	US-11-050-346-45	4
U	20	41.2	7.6	1898	ដ	US-10-450-763-21404	2
	21	36.4	6.7	671	ø	US-10-184-644-346	Sequence 346, App
	22	36.4	6.7	671	9	US-10-184-634-346	Sequence 346, App
O	23	36.2	9.9	3635	m	US-09-312-762A-6	Sequence 6, Appli
O	24	36.2	9.9	3635	2	US-10-764-420-19	Sequence 19, Appl
υ	25	36.2	9.9	3635	ដ	US-10-631-467-1108	Sequence 1108, Ap
υ	56	36	9.9	614	4	US-09-925-065A-434607	Sequence 434607,
υ	27	36	9.9	614	Ŋ	US-09-925-065A-434607	Sequence 434607,
	28	36	9.9	160274	10	US-10-893-315-140	Sequence 140, App
	53	36	9.9	160300	10	US-10-893-315-151	151,
U	30	35.6	6.5	614	4	US-09-925-065A-434608	434606
υ	31	35.6	6.5	614	Ŋ	US-09-925-065A-434608	
O	32	35.6	6.5	919	4	US-09-925-065A-322979	
U	33	35.6	9.5	919	Ŋ	US-09-925-065A-322979	
O	34	35.6	6.5	619	12	US-10-301-480-397487	o
υ	35	35.6	6.5	619	12	US-10-301-480-1010896	
	36	35.6	6.5	165156	8	US-10-741-601-5668	ŧ٨
	37	35.6	6.5	165156	2	US-10-995-561-13304	Sequence 13304, A
υ	38	35.4	6.5	574	12	US-10-301-480-42945	42945,
υ	39	35.4	6.5	574	12	US-10-301-480-656354	Sequence 656354,
U	40	35.4	6.5	585	9	US-10-027-632-181875	Sequence 181875,
U	41	35.4	6.5	585	7	US-10-027-632-181875	Sequence 181875,
v	42	35.4	6.5	4786	13	US-11-097-143-41443	Seguence 41443, A
	43	35.2	9.5	351	m	US-09-867-701-598	98
	44	35.2	9.5	594	9	US-10-123-155-10	10.
	45	35.2	6.5	594	7	-731-1	Sequence 10, Appl
						ALIGNMENTS	
RESULT	17.						
ָרְאָרְ מי	249-01	US-10-698-160-2	יייין	for 11s/	1069	18160	
Š	, i [d	Publication No. HS2004017115941	1320	0401711	5 9 5		
	NERAL	GENERAL INFORMATION	ATION:				

# STREAL INFORMATION: APPLICANT: DC YU APPLICANT: DC YU APPLICANT: DC YU APPLICANT: DC YU APPLICANT: MGGAZAJAN RAMESh TITLE OF INVENTION: CELL-SPECIFIC ADENOVIRUS VECTOR TITLE OF INVENTION: CENPRISING EBV-SPECIFIC PROMOTER FILE REPERBURCE: CELL-0.3 CURRENT APPLICATION NUMBER: US/10/698,160 CURRENT APPLICATION NUMBER: 10/423,203 PRIOR APPLICATION NUMBER: 60/423,203 PRIOR PILING DATE: 2002-11-01 NUMBER OF SEQ ID NOS: 4 SOFTWARE: FASTERQ for Windows Version 4.0 ; TYPE: DNA ; ORGANISM: Epstein Barr virus US-10-698-160-2 LENGTH: 545

ö

Gaps

ö

Length 545;

100.0%; Score 545; DB 8; Length 5: 100.0%; Pred. No. 2.6e-178; Live 0; Mismatches 0; Indels

Query Match Best Local Similarity 100. Matches 545; Conservative

60

120

61 retreaccretercarrectascererecerecerecerecerecerecerecerasaaarea 120

19

8

g à g

g

ઢ

1 CGCACCAACTGCGGCAAATGGCGGTGTTATGAAGGAAAAGGATGGCAGCCTCTCTTTTGC 60 TGTTGACCTGTCACTTCCCAAGCCCTGGCACCTGCCAGTGACCTGCGTTGGGAAAATGA

1 CGCACCAACTGCGGCAAATGGCGGTGTTATGAAGGAAAAGGATGGGAGCCTCTCTGTTGC

CAAGGAGGAAGCCCACGGGGTTTATGTTTTCTGGATACTTGTCGCAATAAACGCACTTGCC 180

121 181

ò

121 CAAGGAGGAAGCCCACGGGGTTTATGTTTCTGGATACTTGTCGCAATAAACGCACTTGCC

| TATTTCACCTTGTTTTAGTGTGGCATTGGGGGGTGGCATTGCGGGTGGATAGCCTGGCG

180

240

```
GenCore version 5.1.9
Copyright (c) 1993 - 2006 Biocceleration Ltd.
```

OM nucleic - nucleic search, using sw model

June Run on:

6, 2006, 18:40:59 ; Search time 36.0089 Seconds (without alignments) 1782.525 Million cell updates/sec

US-10-698-160-2 545

Perfect score:

1 cgcaccaactgcggcaaatg......satttcaggccggttttgca 545 Sequence:

IDENTITY NUC Gapop 10.0 , Gapext 1.0 Scoring table:

Total number of hits satisfying chosen parameters: 246837 segs, 58886990 residues Searched:

seq length: 0 seq length: 200000000 Minimum DB ( Maximum DB (

Post-processing: Minimum Match 0% Maximum Match 100% Listing first 45 summaries

Database :

Published Applications NA New:\*

1: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US09\_NEW\_PUB.seq:\*

2: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US06 NEW\_PUB.seq:\*

3: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US07 NEW\_PUB.seq:\*

4: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US08\_NEW\_PUB.seq:\*

5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US08\_NEW\_PUB.seq:\*

5: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US10\_NEW\_PUB.seq:\*

7: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US11\_NEW\_PUB.seq:\*

7: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US11\_NEW\_PUB.seq:\*

7: /EMC\_Celerra\_SIDS3/ptodata/2/pubpna/US11\_NEW\_PUB.seq:\*

Pred. No. is the number of results predicted by chance to have a score greater than or equal to the score of the result being printed, and is derived by analysis of the total score distribution.

### SUMMARIES

Sequence 3, Appli Sequence 155, App Sequence 655, App Sequence 655, App Sequence 31, Appl Sequence 75811, Ap Sequence 75811, Application	Sequence 6, Appli Sequence 27966, A Sequence 1106, Ap	Sequence 515, App Sequence 136, App Sequence 26032, App Sequence 26032, A	Sequence 1776, Ap Sequence 28, Appl Sequence 1473, Ap Sequence 4118, Ap
US-10-506-549-3 US-11-301-554-155 US-10-953-349-293 US-10-505-928-655 US-11-217-529-2025 US-11-217-529-75811 US-11-269-117-1	US-11-269-117-6 US-10-953-349-27966 US-11-293-697-1106 US-10-196-749-517	US-10-505-928-575 US-11-217-529-136 US-11-953-349-26032 US-11-293-697-367	US-11-293-697-1776 US-10-857-260-28 US-11-301-554-1473 US-10-953-349-4118
91999111	r 9 r 4	96766	r 9 r 9
394191 674 1173 3705 38187 891 1119	8322 1459 2442	5047 1233 1467 1890	3194 16009 526 1544
00000000	0000		0,0,00
01 01 01 01 01 01 01 01	01444	. 4 4 4 4	4444
22222 422 422	26.8 26.8 26.8	26.6 26.6 26.6 26.6	26.6 26.6 26.6 26.4 4.4
33310387	4 5 9 5	8694	4 4 4 4 6 6 6 7
υυ	ט ט	000	υυυ

## ALIGNMENTS

```
571 CTICATCAGCIGIGICACAGGCCAAAGAACCIAGTIGIGCCACTICAAAAGGAIGIAACC 512
            Sequence 77277, Application US/11217529

Publication No. US20060099612A1

GENERAL INFORMATION:

APPLICANT: SUNTORY LIMITED

APPLICANT: NAKAMURA, VORHIERO

APPLICANT: NAKAMURA, TOWOKO

APPLICANT: KUDAMA, YUKIKO

APPLICANT: RUJIMURA, TOWOKO

APPLICANT: ASHIKARI, TOSHIHIKO

APPLICANT: ASHIKARI, TOSHIHIKO

TITLE OF INVENTION: METHODS FOR ANALYZING GENES OF INDUSTRIAL YEASTS

FILE REPERENCE: S-38-285

CURRENT APPLICATION NUMBER: US/11/217,529

FILE REPERENCE: 2005-09-02

PRIOR FILING DATE: 2005-09-02

PRIOR FILING DATE: 2004-09-02

NUMBER OF SEQ ID NOS: 197023

SOFTWARE: PATENTIN VERSION 3.3
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           74 CTTCCCAAGCCCTGGCACCTGCCAGTGACCTGCGTTGGGAAAAATGACAAGGAGGAAGCC
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 Gaps
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                 .
0
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  Length 666;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          39; Indels
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                          5.6%; Score 30.6; DB 7;
58.1%; Pred. No. 0.85;
iive 0; Mismatches 39;
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     134 CACGGGGTTTATGTTTCTGGATACTTGTCGCAA 166
                                                                                                                                                                                                                                                                                                                                                                                                                                                                               ; TYPE: DNA
; ORGANISM: Saccharomyces pastorianus
US-11-217-529-77277
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        Query Match
Best Local Similarity 58.11
Matches 54; Conservative
US-11-217-529-77277/c
                                                                                                                                                                                                                                                                                                                                                                                                                                  SEQ ID NO 77277
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                           ò
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                        g
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                       ઠે
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                         g
```

ö

Sequence 382, Application US/11293697
Publication No. US20060105376A1
GENERAL INFORMATION:
TITLE OF INVENTION: Novel full length CDNA
FILE REPRENCE: HI-A0106
CURRENT APPLICATION WURBER: US/11/293,697
CURRENT FILING DATE: 2005-12-05 RESULT 2 US-11-293-697-382/c

#### FOR OFFICIAL USE ONLY

191632 PLEASE PRINT CLEARLY

Scientific and Technical Information Center

#### SEARCH REQUEST FORM

(54.0)	
Requester's Full Name: DAVID GUZO Examiner #: 70677 Date: 6/1/06	
Requester's Full Name: DAVID GUZO Examiner #: 70677 Date: 6/1/06  Art Unit: 1636 Phone Number: 2-0767 Serial Number: 10/698160	
Location (Bldg/Room#): 2070 Results Format Preferred (circle): PAPER DISK	
To ensure an efficient and quality search, please attach a copy of the cover sheet, claims, and abstract or fill out the following:	
Title of Invention:	
Inventors (please provide full names):	
Earliest Priority Date:	
Search Topic:  Please provide a detailed statement of the search topic, and describe as specifically as possible the subject matter to be searched. Include th elected species or structures, keywords, synonyms, acronyms, and registry numbers, and combine with the concept or utility of the invention Define any terms that may have a special meaning. Give examples or relevant citations, authors, etc., if known.	
*For Sequence Searches Only* Please include all pertinent information (parent, child, divisional, or issued patent numbers) along with the appropriate serial number.	e
Please una regular plus interferens seguerce	

search on SEQ IO NO: 1 and 2.

1-575 Thanks 3-545